

## UNITED STATES PATENT AND TRADEMARK OFFICE

UNITED STATES DEPARTMENT OF COMMERCE United States Patent and Trademark Office Address: COMMISSIONER FOR PATENTS P.O. Box 1450 Alexandria, Virginia 22313-1450 www.uspto.gov

CONFIRMATION NO. FIRST NAMED INVENTOR ATTORNEY DOCKET NO. FILING DATE APPLICATION NO. 4856 1064/48929 Markku Michael Jeltsch 07/26/2001 09/912,436 **EXAMINER** 11/09/2004 23911 SPECTOR, LORRAINE CROWELL & MORING LLP INTELLECTUAL PROPERTY GROUP PAPER NUMBER ART UNIT P.O. BOX 14300 1647 WASHINGTON, DC 20044-4300

DATE MAILED: 11/09/2004

Please find below and/or attached an Office communication concerning this application or proceeding.



# United States Patent and Trademark Office



UNITED STATES DEPARTMENT OF COMMERCE United States Patent and Trademark Office Address: COMMISSIONER FOR PATENTS P.O. Box 1450 Alcandria, Virgaia 22313-1450 www.uspia.gov

APPLICATION NO.	FILING DATE	FIRST NAMED INVENTOR	ATTORNEY DOCKET NO.	CONFIRMATION NO.
09/912,436	07/26/2001	Markku Michael Jeltsch	1064/48929	4856
23911	7590 01/09/2004		EXAMINER	
CROWELL & MORING LLP			SPECTOR, LORRAINE	
INTELLECTUAL PROPERTY GROUP			ART UNIT	PAPER NUMBER
P.O. BOX 14300 WASHINGTON; DC 20044-4300			1647	6
	3		DATE MAILED: 01/09/200	4 16

Please find below and/or attached an Office communication concerning this application or proceeding.

·	I A a strategy No.	Applicant(s)				
	Application No.	JELTSCH ET AL.				
Office Action Summary	09/912,436 Examiner	Art Unit				
Omce Housen Cummary	Lorraine Spector, Ph.D.	1647				
The MAILING DATE of this communication ap						
Period for Reply		•				
A SHORTENED STATUTORY PERIOD FOR REPL THE MAILING DATE OF THIS COMMUNICATION.  - Extensions of time may be available under the provisions of 37 CFR 1.  - after SIX (6) MONTHS from the mailing date of this communication.  - If the period for reply specified above is less than thirty (30) days, a rep  - If NO period for reply is specified above, the maximum statutory period  - Failure to reply within the set or extended period for reply will, by statut  - Any reply received by the Office later than three months after the mailin  - earned patent term adjustment. See 37 CFR 1.704(b).	136(a). In no event, however, may a reply be till ly within the statutory minimum of thirty (30) da will apply and will expire SIX (6) MONTHS from a cause the application to become ABANDONI	mely filed ys will be considered timely. In the mailing date of this communication. ED (35 U.S.C. § 133).				
Status						
1) Responsive to communication(s) filed on						
24)						
3) Since this application is in condition for allowance except for formal matters, prosecution as to the merits is closed in accordance with the practice under Ex parte Quayle, 1935 C.D. 11, 453 O.G. 213.						
Disposition of Claims						
4) Claim(s) 1-18 is/are pending in the application.						
4a) Of the above claim(s) is/are withdra	4a) Of the above claim(s) is/are withdrawn from consideration.					
5) Claim(s) is/are allowed.	5) Claim(s) is/are allowed.					
6)⊠ Claim(s) <u>1-18</u> is/are rejected.						
	Claim(s) is/are objected to.					
8) Claim(s) are subject to restriction and/o	or election requirement.					
Application Papers						
9) The specification is objected to by the Examiner.						
	) ☐ The drawing(s) filed on is/are: a) ☐ accepted or b) ☐ objected to by the Examiner.					
	Applicant may not request that any objection to the drawing(s) be held in abeyance. See 37 CFR 1.85(a).					
Replacement drawing sheet(s) including the correction is required if the drawing(s) is objected to. See 37 CFR 1.121(d).						
11) The oath or declaration is objected to by the E	xaminer. Note the attached Office	e Action or form P1O-152.				
Priority under 35 U.S.C. §§ 119 and 120						
12) Acknowledgment is made of a claim for foreign priority under 35 U.S.C. § 119(a)-(d) or (f).  a) All b) Some * c) None of:  1. Certified copies of the priority documents have been received.  2. Certified copies of the priority documents have been received in Application No.  3. Copies of the certified copies of the priority documents have been received in this National Stage application from the International Bureau (PCT Rule 17.2(a)).  * See the attached detailed Office action for a list of the certified copies not received.  13) Acknowledgment is made of a claim for domestic priority under 35 U.S.C. § 119(e) (to a provisional application) since a specific reference was included in the first sentence of the specification or in an Application Data Sheet.  37 CFR 1.78.  a) The translation of the foreign language provisional application has been received.  14) Acknowledgment is made of a claim for domestic priority under 35 U.S.C. §§ 120 and/or 121 since a specific reference was included in the first sentence of the specification Data Sheet. 37 CFR 1.78.						
Attachment(s)	4) Then iew Summar	y (PTO-413) Paper No(s)				
1) Notice of References Cited (PTO-892) 2) Notice of Draftsperson's Patent Drawing Review (PTO-948)	5) Notice of Informal	Patent Application (PTO-152)				
3) Information Disclosure Statement(s) (PTO-1449) Paper No(s)	<u>5/15/02</u> . 6) ☐ Other: .					

Application/Control Number: 09/912,436

Art Unit: 1647

#### Part III: Detailed Office Action

Claims 1-18 are pending and under consideration.

#### Formal Matters:

The disclosure is objected to because it contains an embedded hyperlink and/or other form of browser-executable code. For example, see page 13. Applicant is required to delete the embedded hyperature and/or other form of browser-executable code. See MPEP § 608.01.

### Objections and Actions under 35 U.S.C. \$112:

The following that tation of the second paragraph of 35 U.S.C. 112:

The specification shall conclude with one or more claims particularly pointing out and distinctly claiming the subject matter which the applicant regards as his invention.

Claims 1-13 are rejected under 35 U.S.C. 112, second paragraph, as being indefinite for failing to particularly point out and distinctly claim the subject matter which applicant regards as the invention.

The circle indefinite because they both require a particular sequence, and require that that sequence deed, that is, not be the recited sequence. Using claim 1 as an example, the claim has a second that may be SEQ ID NO: 1, 3 or 5, and "a nucleotide sequence encoding at least one putal description of site inserted therein", so that it would no longer be SEQ ID NO: 1, 3 or 5. The age such as "consisting of a variant of SEQ ID NO: 1, 3 or 5 which consists of SEQ ID NO: 3 which has been altered to add at least one putative N-glycosylation site" or the equival and be remedial.

The class also indefinite because of the recitation that the glycosylation site has been "inserted the protein. Art-accepted usage of the word "inserted" or "insertion" denotes a length of substitution that the glycosylation as disclosed is the result of substitution acids to form a glycosylation site without lengthening the resultant molecule. We goal can also be achieved by insertion of additional codons, such would seem to be I the invention to exclude the working example. As it is not clear what applicants into the serting, the claims that recite such are indefinite.

Application/Con Number: 09/912,436

Art Unit: 1647

Claim 1 ther indefinite because the metes and bounds of those nucleic acids that will hybridize to a sequence are dependent upon the conditions used for hybridization and washing. The "stringent conditions" is a relative term, and neither the claim nor the specification broad life and meaning into the term to allow determination of the metes and bounds of the conditions. It is noted that exemplary hybridization conditions are disclosed at page 13 of the specific and, however said definition is incomplete, and is merely exemplary, and non-limiting.

Claim 1 definite because it is not clear what is intended by "increasing an amount of a soluble VEC." from a host cell"; amendment to replace "from" with "secreted by " or "produced by" be remedial.

The re. sclaims are rejected for depending from an indefinite claim.

The following is a quotation of the first paragraph of 35 U.S.C. 112:

The species of collider contain a written description of the invention, and of the manner and process of making and using it, in such dill, clear, concise, and exact terms as to enable any person skilled in the art to which it pertains, or with which is structured as a concept, to make and use the same and shall set forth the best mode contemplated by the inventor contemplated by the inventor.

Claims 3, 5, 7, and 8 are rejected under 35 U.S.C. 112, first paragraph, as containing subject matter which was not described in the specification in such a way as to reasonably convey to one illed to the relevant art that the inventor(s), at the time the application was filed, had possessic fithe sammed invention.

The claims are drawn to nucleic acids of unspecified structure. Claim 1 requires only that the claimed nucleic acid "hybridizes under stringent conditions" to a reference sequence, and have "at least the putative N-glycosylation site inserted therein". It is generally accepted in the art that only a limited stretch of identity is required for two sequences to hybridize under "stringent" conserved structural feature that would lend function to the encoded any significant conserved structural feature that would lend function to the encoded protein. Further, the coloms have no functional limitations as to either the nucleic acid itself of any protein that it. In the encoded thereby. The specification, on the other hand, is clearly drawn to nucleic acids to the encoded thereby. VEGF-B, the sequence of which has been altered to

comprise one or more putative N-linked glycosylation sites, which sites are characterized by the amino acid sequence "NXT". There is no evidence of conception of an invention commensurate with the breadth being claimed.

Vas-Cath Inc. v. Mahurkar, 19USPQ2d 1111, clearly states that "applicant must convey with reasonable clarity to those skilled in the art that, as of the filing date sought, he or she was in possession of the invention. The invention is, for purposes of the 'written description' inquiry, whatever is now claimed." (See page 1117.) The specification does not "clearly allow persons of ordinary skill in the art to recognize that [he or she] invented what is claimed." (See Vas-Cath at page 1116).

With the exception of the sequences referred to above, the skilled artisan cannot envision the detailed chemical structure of the encompassed polynucleotides, and therefore conception is not achieved until reduction to practice has occurred, regardless of the complexity or simplicity of the method of isolation. Adequate written description requires more than a mere statement that it is part of the invention and reference to a potential method of isolating it. The nucleic acid itself is required. See Fiers v. Revel. 25 USPQ2d 1601 at 1606 (CAFC 1993) and Amgen Inc. v. Chugai Pharmaceutical Co. Ltd., 18 USPQ2d 1016.

One cannot describe what one has not conceived. See *Fiddes v. Baird*, 30 USPQ2d 1481 at 1483. In *Fiddes*, claims directed to mammalian FGF's were found to be unpatentable due to lack of written description for the broad class. The specification provided only the bovine sequence.

Therefore, only SEQ ID N 1, 3 or 5, modified by the introduction of N-glycosylation sites, but not the full breadth of the claim meets the written description provision of 35 U.S.C. §112, first paragraph. Applicant is reminded that *Vas-Cath* makes clear that the written description provision of 35 U.S.C. §112 is severable from its enablement provision (see page 1115).

Claims 1-5, 7-13, and 15 17 are rejected under 35 U.S.C. 112, first paragraph, as containing subject matter which was not described in the specification in such a way as to enable one skilled in the art to which it artains, or with which it is most nearly connected, to make and/or use the invention.

The factors considered when determining if the disclosure satisfies the enablement requirement and whether any necessary experimentation is "undue" include, but are not limited to:

1) nature of the invention, 2) state of the prior art, 3) relative skill of those in the art, 4) level of predictability in the art, 5) existence of working examples, 6) breadth of claims, 7) amount of direction or guidance by the inventor, and 8) quantity of experimentation needed to make or use the invention. In re Wands, 858 F. 11 731, 737, 8 USPQ2d 1400, 1404 (Fed. Cir. 1988).

claims, however, is extreme. The they limited by particular structure of the claims. There is no stre acids, which themselves are not the person of ordinary skill in the VEGF-B and nucleic acids en specification does not provide defined by "hybridization" lan activity, or proteins without Accordingly, it would require u commensurate in scope with the

The nature of the invention is the insertion of N-glycosylation sites in to VEGF-B. The state of the prior art, as discussed below, is that VEGF-B was known, that it was known that the protein was not glycosylated, that it was further known that glycosylation of VEGF-A increases recombinant production of the pull-in without affecting biological activity, and finally, it was known in the art how to introduce a ycosylation sites into proteins, although it is not predictable whether or not a given glycosy. On site will actually be glycosylated. The breadth of the aims are not limited to proteins with VEGF activity, nor are iue to the inclusion of "hybridization" language in numerous onor function to be conserved by the protein or the nucleic aired, as in claim 1, to actually encode any protein. While would certainly know how to make glycosylated variants of ag such, and would further know how to use such, the date guidance as to how to make or use variants that are , nucleic acids that do not encode protein having VEGF 3 function or having distinct but unspecified function. experimentation to make and use the invention in a manner is.

The following is a quota obviousness rejections set forth (a) A patent may not be obtained 102 of this title, if the difference subject matter as a whole would in the art to which said subject it invention was made.

f 35 U.S.C. 103(a) which forms the basis for all

Office action:

the invention is not identically disclosed or described as set forth in section a the subject matter sought to be patented and the prior art are such that the n obvious at the time the invention was made to a person having ordinary skill stains. Patentability shall not be negatived by the manner in which the

This application currently claims under 35 U.S.C. 103(a), claims was commonly owned at evidence to the contrary. Application the inventor and invention dates convention was made in order for and potential 35 U.S.C. 102(e), (C.)

Claims 1-9, 11, and 13-400 Eriksson et al. (U.S. Patent Num. and Keyt et al. (U.S. Patent Num.)

Eriksson et al. disclosconservation among PDGF-A a figure that the sole glycosylation residues 65-67 of VEGF-B as making they are highly similar, and glycosylation site of VEGF-A compositions comprising the VEGF-A

Eriksson et al. do not protein or making nucleic acic compositions comprising hepar

Claffey et al. disclose a Tyrosine, thus eliminating the page 7, they state that the action wild-type, glycosylated form, that of wild-type. Claffey et al.

N-linked glycosylation sites a encode such as site, and that motifs. Addition of glycosylat

examiner presumes that the subject matter of the various time any inventions covered therein were made absent any is advised of the obligation under 37 CFR 1.56 to point out ach claim that was not commonly owned at the time a later examiner to consider the applicability of 35 U.S.C. 103(c) (g) prior art under 35 U.S.C. 103(a).

rejected under 35 U.S.C. 103(a) as being unpatentable over 5,607,918) in view of Claffey et al. (BBA 1246:1-9, 1995) 3,020,473).

3F-B. At figure 9, an alignment is provided, showing 3, PLGF, VEGF, and VEGF-B. It can be seen from that of VEGF-A ("NIT") occurs at a position corresponding to cred in the current specification ("QVR") The VEGF-B of see and human VEGF-B are aligned, and it can be seen that ical in sequence at the position corresponding to the combinant expression of the protein and pharmaceutical 3 protein are claimed.

se introducing added N-glycosylation into the VEGF-B ding such a species, nor do they disclose pharmaceutical VEGF-B.

nt of VEGF-A in which Asn at residue 74 is replaced with linked glycosylation site in the protein - see Figure 3. At the non-glycosylated form was comparable to that of the secretion of the non-glycosylated form was only 50% of

EGF-A. At column 8, they teach that the protein may have it via recombinant expression of nucleic acids altered to ed glycosylation may occur at Asn-X-Ser or Asn-X-Thr s is also discussed at column 13. At columns 35-36, Keyt

Application/Control Number: 057

Art Unit: 1647

discuss the results of glycosylation that elimination of glycosylation showed that glycosylation via interfere with receptor binding binding to the KDR receptor, as but not KDR binding.

It would have been obtainvention was made to modification site at the position. Such introduction of new glycons. Keyt et al., and the person of commodification in view of Clafford better than, and has activity at Eriksson's alignment of the relevance of the relevance of the person of ordinary skill expected the VEGF-B so obtains as well or better than non-glyconselected protein. Accordingly, cited prior art.

Claims 10 and 12 ar Eriksson et al. (U.S. Patent N and Keyt et al. (U.S. Patent N and further in view of Erikss Patent Number 5,851,989).

Claim 10 recites that t recites that the host cell expensed.

added variants of VEGF-A. They reiterate Claffey's result esidue 75 did not affect receptor binding. They additionally action of a "neoglycosylation site" at residues 42-44 did not that glycosylation at residues 82-84 resulted in decreased tra glycosylation at position 64 was shown to decrease Flt-1

to the person of ordinary skill in the art at the time the GF-B, as taught by Eriksson et al., by introducing a sponding to the position at which VEGF-A is glycosylated. On sites is routine in the art, as taught by Claffey et al. and ry skill in the art would have been motivated to make the d Keyt's teachings that glycosylated VEGF-A is secreted ant to, non-glycosylated VEGF-A, and taken in view of oteins, including VEGF-A and VEGF-B, thus pointing out ion is. The possibility of better secretion would be ample the would facilitate recombinant production of the protein art, making this modification, would reasonably have retain its biological function, and to be secreted from cells I VEGF-B, as that was the result with VEGF-A, a closely ention, taken as a whole, is *prima facie* obvious over the

ed under 35 U.S.C. 103(a) as being unpatentable over 6,607,918) in view of Claffey et al. (BBA 1246:1-9, 1995) 6,020,473) as applied to claims 1-9, 11, and 13-18 above, E.S. Patent Number 5,840,693) and Chamow et al. (U.S.

the protein is exposed to heparin after the protein is

Eriksson -2 discloses the release of VEGF-B dimers comprising VEGF-B and heparities.

Chamow et al. disclose a proteins such as VEGF; see ab.

It would have been observation was made to treat obvious above with heparin as secreted dimers, as taught by E

It would have been furtiinvention was made to form
heparin as taught by both Er
pharmaceutical with an extendave expected success, in viteachings of the advantages of subject matter, taken as a who

atment of cells expression VEGF-B with heparin results in the cells, see col. 20, and additionally claims compositions claims 10 and 11.

heparin increases the serum half-life of heparin binding nd claims.

to the person of ordinary skill in the art at the time the expressing recombinant glycosylated VEGF-B as found t by Eriksson-2, for the purpose of increasing the yield of -2.

rious to the person of ordinary skill in the art at the time the compositions comprising the glycosylated VEGF-B and and by Chamow et al., for the purpose of obtaining a life. One would have been motivated to do so, and would be claims of both of the secondary references, and the found in the Chamow patent. Accordingly, the claimed ha facie obvious over the prior art.

#### Conclusion

No claim is allowed.

Any inquiry concerning Examiner should be directed 1793. Dr. Spector can normal Effective 1/21/2004, Dr. Spec

If attempts to reach supervisor, Dr. Gary L. Kunnumber will be 571-272-088?

Any inquiry of a gene should be directed to the Grov

Certain papers related transmission The faxing of some Gazette, 1156 OG 61 (Novel C.F.R. § 1.6(d)). NOTE: It should be retained by applications of the SHOULD BE SUBMITTED.

communication or earlier communications from the ine M. Spector, whose telephone number is (703) 308-eached Monday through Friday, 9:00 A.M. to 5:30 P.M. phone number will be 571-272-0893.

miner by telephone are unsuccessful, the Examiner's 3)308-4623. Effective 1/21/2004, Dr. Kunz' telephone

or relating to the status of this application or proceeding nist at telephone number (703) 308-0196.

pplication may be submitted to Group 1800 by facsimile must conform with the notices published in the Official 1993) and 1157 OG 94 (December 28, 1993) (see 37 t does submit a paper by fax, the original signed copy pplicant's representative. NO DUPLICATE COPIES bid the processing of duplicate papers in the Office.

Official papers filed to (703)872-9307 (after finashould be directed to (703) 7 571-273-0893.

ould be directed to (703) 872-9306 (before final rejection) duraft or informal communications with the examiner Effective 1/21/2004, Dr. Spector's fax number will be

Lorraine Spector, Ph.D. Primary Examiner